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Rhodium-Catalyzed Borylation of Aryl 2-Pyridyl Ethers through Cleavage of the Carbon-Oxygen Bond: Borylative Removal of the Directing Group

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ABSTRACT: The rhodium-catalyzed reaction of aryl 2-pyridyl ethers with a diboron reagent results in the formation of arylboronic acid derivatives via activation of the C(aryl)-O bonds. The straightforward synthesis of 1,2-disubstituted arenes was enabled through catalytic ortho C-H bond functionalization directed by the 2-pyridyloxy group followed by substitution of this group with a boryl group. Several control experiments revealed that the presence of a sp² nitrogen atom at the 2-position of the substrate and the use of a boron-based reagent were crucial for the activation of the relatively inert C(aryl)-O bond of aryl 2-pyridyl ethers.

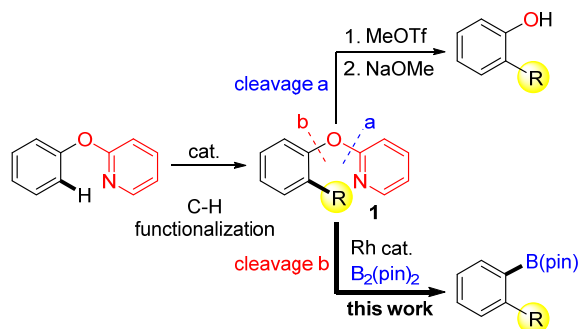
Introduction

Transition metal-catalyzed transformation of C-H bonds has emerged as a powerful synthetic method, because it eliminates the need for pre-activation of the starting materials.^{1,2} Although the ubiquitous nature of C-H bonds makes this strategy highly versatile, this ubiquity results in a regioselectivity issue: which C-H bond reacts. Although the regioselectivity has often been controlled by the differentiation in the electronic and steric nature of the C-H bonds, these factors are substrate-dependent.¹ An alternative way to realize regioselective C-H functionalization involves the use of a directing group.^{2,3} Metal-coordinating groups, such as ketones, amides, esters, nitriles, and alcohols, have been used as directing groups, which control the regioselectivity by forming stable metallacyclic intermediates. Among these metal-coordinating groups, the pyridine ring is a frequently used motif, and 2-phenylpyridine is a privileged substrate for investigating new ortho C-H bond functionalization reactions.^{4,5} The utility of the 2-pyridyl moiety is attributable in part to the strong coordination ability of the sp² nitrogen atom and its stability under various transition metal-catalyzed conditions. Despite the outstanding performance of 2-phenylpyridine substrates in C-H bond functionalization reactions, the utility of the products is severely limited because it is nontrivial to remove and functionalize the 2-pyridyl moiety. To overcome this limitation, several modified 2-pyridyl groups have been de-

veloped.⁶ The 2-pyridylsilyl group is a suitable directing group in several catalytic ortho C-H transformations, and can subsequently be converted to a reactive group, such as a halide.⁷ However, introduction of this useful directing group requires the tedious preparation of organosilicon compounds. 2-Aminopyridine can also serve as a removable directing group,⁸⁻¹⁰ although the aniline product requires activation, such as diazotization, for further elaboration. Perhaps the most useful metal-coordinating group is 2-pyridyloxy (OPy), which serves as an excellent ortho directing group in a number of C-H bond functionalization reactions¹¹ and can subsequently be removed. The only available method to remove the pyridine ring in 2-OPy involves (1) *N*-methylation and (2) cleavage of the C(pyridinium)-O bond by NaOMe to give the corresponding phenol (Scheme 1, cleavage a).^{11c} Although the development of this method has significantly increased the synthetic utility of the OPy directing group, several aspects of the method require improvement: (1) the use of a strong methylating reagent and a strong base limits applicable functional groups, (2) the protocol for the removal of the pyridine ring requires two steps, and (3) an additional step is required for further elaboration of phenols (e.g., conversion to triflates). In this Article, we report a new method for the conversion of the OPy group by substituting the OPy group with a synthetically useful boryl group in a single step and in a catalytic manner (Scheme 1, cleavage b).¹² In this reaction, the relatively electron-rich C(aryl)-O bond in ether **1** is selectively cleaved over the

electron-deficient C(pyridyl)-O bond.^{13,14} This borylation proceeds under neutral conditions and is operationally simple.

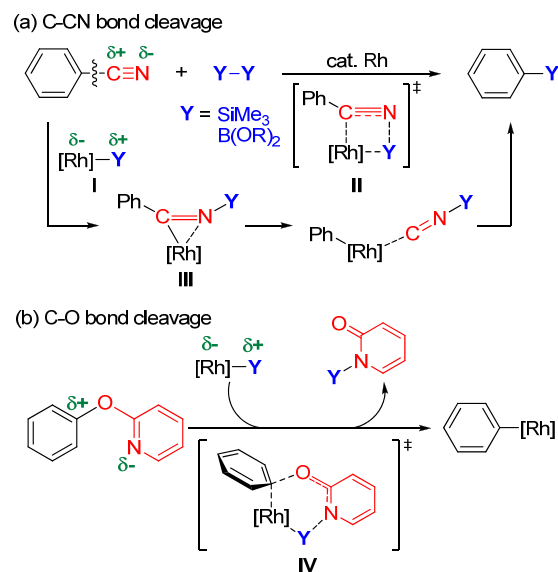
Scheme 1. 2-Pyridyloxy Group as an Ortho Directing Group in C-H Bond Functionalization and Its Subsequent Derivatization



Results and Discussion

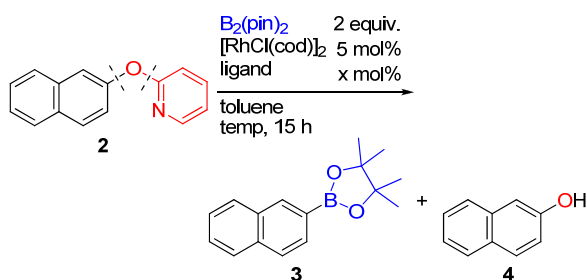
Previously, we developed a series of rhodium-catalyzed transformation reactions of nitriles through the cleavage of the C-CN bonds in the presence of organosilicon¹⁵ or diboron¹⁶ reagents. Stoichiometric¹⁷ and theoretical studies^{18,19} revealed that the silylrhodium or borylrhodium species I (depicted as [Rh]-Y) generated in situ mediates the cleavage of the C-CN bond through iminoacyl intermediate III (Scheme 2a).²⁰ The key feature of catalytically active species I is the Lewis acid nature of the silyl^{21,22} or boryl ligand,^{23,24} which favors binding to the Lewis base nitrogen of the cyano group in transition state II. This polarity-driven interaction allows for the formation of iminoacyl intermediate III, which eventually results in the cleavage of the carbon-carbon bond. These mechanistic considerations led us to hypothesize that the unique polarity of I could be applied to the activation of the C-OPy bond via cyclic transition state IV, in which the boron-nitrogen interaction facilitates the otherwise difficult C-O bond cleavage (Scheme 2b).

Scheme 2. Mechanism of Silylrhodium- or Borylrhodium-Mediated C-CN Bond Cleavage and Working Hypothesis for Application to C-O Bond Cleavage

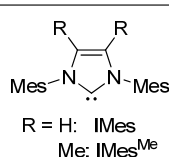
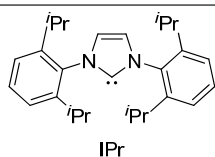


We began our study by investigating the reaction of aryl pyridyl ether **2**, which was easily prepared from 2-naphthol and 2-bromopyridine, with bis(pinacolato)diboron ($B_2(\text{pin})_2$) in the presence of a rhodium(I) catalyst (Table 1). Pleasingly, it was found that the use of the rhodium(I) catalyst in conjunction with PPh_3 gave arylboronate **3** via the cleavage of the $C(\text{sp}^2)\text{-O}$ bond (Table 1, entry 1). Considering the widespread use of the OPy group as a directing group in C-H bond activation reactions,¹¹ it is notable that the ortho C-H bonds remained unchanged under these conditions. However, the conversion of **2** was low (54%) and the selectivity of the cleavage of the two C-O bonds was moderate (3:4 = 3:1). Attempts to improve the reactivity and selectivity by introducing electron-donating or -withdrawing groups on the pyridine ring and by replacing the 2-pyridyl moiety with other nitrogen-containing heteroaromatics were unsuccessful.²⁵

Next, we turned our attention to the effect of the ligand. It was found that the use of a more electron-rich phosphine ligand was better in terms of both conversion and selectivity (Table 1, entry 2). The selectivity dramatically decreased when bis(neopentylglycolato)diboron was used in place of $B_2(\text{pin})_2$ (entry 3). When the reaction was performed using PCy_3 as the ligand, the highest conversion was achieved and **3** was obtained as the only product (entry 4). Further improvement in the yield was achieved by decreasing the temperature to 100 °C, with **3** being formed in 89% yield (entry 5). The use of *N*-heterocyclic carbene (NHC) ligands, such as IMes, was also effective for selectively forming **3**, although the yield was slightly lower than that obtained with PCy_3 (entry 8).

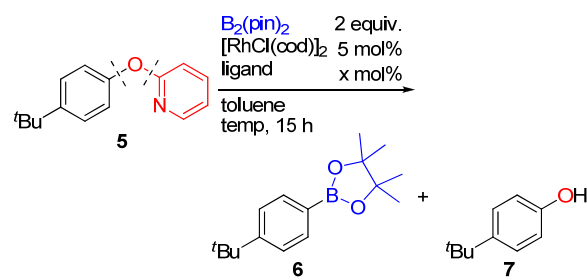
Table 1. Optimization Study of the Borylation of 2^a

entry	ligand (mol%)	temp (°C)	NMR yields (%)		
			3	4	2
1	PPh ₃ (30)	130	26	9	46
2	P(4-MeOC ₆ H ₄) ₃ (30)	130	35	9	37
3 ^b	P(4-MeOC ₆ H ₄) ₃ (30)	130	30	35	16
4	PCy ₃ (30)	130	65	0	0
5	PCy₃ (30)	100	89	0	0
6	PCy ₃ (30)	80	4	0	89
7	IPr (20)	130	33	2	4
8	IMes (20)	130	70	0	0
9	IMes (10)	130	36	0	0
10	IMes (20)	100	27	0	72
11	IMes ^{Me} (20)	130	59	0	0

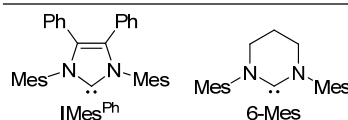


^a Reaction conditions: 2 (0.50 mmol), B₂(pin)₂ (1.0 mmol), [RhCl(cod)]₂ (0.025 mmol), ligand, toluene (0.50 mL) for 15 h.
^b Bis(neopentylglycolato)diboron was used instead of B₂(pin)₂.

Having identified PCy₃ as the optimal ligand for the rhodium-catalyzed borylative cleavage reaction of 2, we next applied the optimized protocol to other substrates. However, unfortunately, the optimized conditions in Table 1 were ineffective for several other less reactive substrates. For example, the borylation of pyridyl ether 5 using the PCy₃ ligand did not give borylated product 6 (Table 2, entry 1). This prompted us to reinvestigate the reaction conditions for less reactive substrates, such as 5. Inspired by the result that the use of IMes gave the borylated product in 22% yield (entry 3), we focused on modification of the IMes ligand. Introduction of substituents at the 4- and 5- positions of the imidazolide core of IMes dramatically improved the yield of 6 (entries 5-8). Among the substituted IMes ligands, IMes^{Me} was found to be optimal, with 6 being obtained in 77% yield with no 7 being generated (entry 5). A similar result was also obtained when [RhCl(C₂H₄)₂]₂ was used as the catalyst precursor (entry 6). Other NHC ligands containing a six-membered framework were ineffective for this borylation reaction (entries 9 and 10).

Table 2. Optimization Study of the Borylation of 5^a

entry	ligand (mol%)	temp (°C)	NMR yields (%)		
			6	7	5
1	PCy ₃ (30)	100	0	38	21
2	PCy ₃ (30)	130	18	34	30
3	IMes (20)	130	22	3	7
4	IMes (20)	100	0	0	99
5	IMes^{Me} (20)	130	77	0	0
6 ^b	IMes ^{Me} (20)	130	76	0	0
7	IMes ^{Me} (20)	100	31	0	54
8	IMes ^{Ph} (20)	130	56	0	0
9	6-Mes (20)	130	15	4	1
10	6-Mes (20)	80	13	2	15



^a Reaction conditions: 5 (0.50 mmol), B₂(pin)₂ (1.0 mmol), [RhCl(cod)]₂ (0.025 mmol), ligand, toluene (0.50 mL) for 15 h.
^b [RhCl(C₂H₄)₂]₂ was used as the Rh source.

With the two different sets of conditions (condition A: PCy₃, 100 °C; condition B: IMes^{Me}, 130 °C), we then investigated the scope of the rhodium-catalyzed borylation reaction using a variety of aryl pyridyl ether substrates (Table 3). Condition B was successful for pyridyl ethers containing a range of functional groups, including simple ethers (entries 3, 4, and 22), fluorinated substituents (entries 5, 6, and 9), and amines (entry 17). It should be noted that this catalytic system exhibited excellent selectivity for reactions containing different C-O bonds. The Ar-OPy bond was exclusively borylated with the Ar-OMe (entry 3), Ar-OPh (entry 4), Ar-OPiv (entry 13) and Ar-OCONMe₂ (entry 15)²⁷ groups remaining completely intact under these catalytic conditions, although all of the latter C-O bonds have been reported to be reactive toward nickel-catalyzed reactions.²⁸ Condition B proved to be unsuitable for substrates containing carbonyl functionalities (entries 12, 14, 16, and 36). This issue was addressed by using condition A, which allows for the borylation of pyridyl ethers containing esters (entries 11 and 13), carbamates (entries 15 and 34), and amides (entry 35). This borylation reaction was found to be relatively sensitive to steric effects: ortho substituted substrates gave the corresponding products in relatively lower yields (entries 18 and 20). The decrease in the yields of the borylated products was found to be due

1
2
3
4
5 to the formation of a reductive cleavage product.²⁵ These
6 results indicate that an ortho substituent does not signifi-
7 cantly inhibit the C(aryl)-O activation process but rather
8 retards the subsequent oxidative addition of B₂(pin)₂ (D
9 → F in Scheme 4 below). To accelerate this step, we de-
10 signed a new NHC ligand that should serve as a stronger
11 σ-donor than IMes^{Me}. As a result, a methoxy-substituted
12 analogue IMXy^{Me} was found to be a better ligand for or-
13 thio-substituted substrates.²⁵ Under the reoptimized con-
14 ditions using IMXy^{Me} (condition C), the yields from sub-
15 strates **19** and **20** were increased to 63% (entry 19) and
16 70% (entry 21), respectively. Although ketones and alde-
17 hydes were incompatible in this catalytic system, the use
18 of an acetal protecting group (entry 25) allows application
19 to such substrates. A range of heteroaromatic substrates,
20 including quinoline (entry 28), indole (entry 31), carbazole
21 (entry 32), and thiophene (entry 33) successfully under-
22 went the borylation reaction. This protocol can also be
23 applied to complex molecules derived from tyrosine (en-
24 try 34) and proline (entry 35). In addition, the C(sp₃)-OPy
25 bond in benzyl alcohol derivatives can be activated in this
26 Rh/diboron system (entries 37 and 39).

27
28 (Table 3 can be found in page 9)

29
30 Because aryl 2-pyridyl ethers can be readily obtained
31 from the corresponding phenol and 2-bromopyridine, and
32 can readily undergo a number of ortho C-H bond func-
33 tionalization reactions, the present borylation should
34 provide a new strategy for the synthesis of a range of 1,2-
35 disubstituted arenes from simple phenol derivatives
36 (Scheme 3). For example, the ruthenium-catalyzed reac-
37 tion of **15** with ethyl acrylate¹⁸ followed by hydrogenation
38 gave ortho alkylated product **33**, which can be borylated
39 via the loss of a OPy group to form arylboronate **34**. Simi-
40 larly, introduction of an ortho aryl group via ruthenium
41 catalysis^{18d} followed by borylation via rhodium catalysis
42 allows for the straightforward synthesis of 2-
43 arylphenylboronic acid derivatives starting from phenol.
44 The OPy directing group can also promote ortho alkoxy-
45 lation by palladium catalysis to form **37**,^{18k} which eventual-
46 ly leads to the borylated product **38** using our method.
47 When the palladium-catalyzed reaction was performed in
48 a carbon monoxide atmosphere, an ester group could also
49 be installed at the ortho position to form **39**.^{18j} The OPy
50 moiety was again converted to the boryl group under the
51 same rhodium-catalyzed conditions to give **40**. Thus,
52 highly functionalized arylboronates, such as **34**, **36**, **38**,
53 and **40**, can be synthesized and serve as useful building
54 blocks that are amenable to further elaboration.²⁹ For
55 example, compound **40**, which contains an ortho ester
56 group, can be annulated with alkyne to form indenone
57 derivative **41** in a single step under palladium-catalysis.³⁰

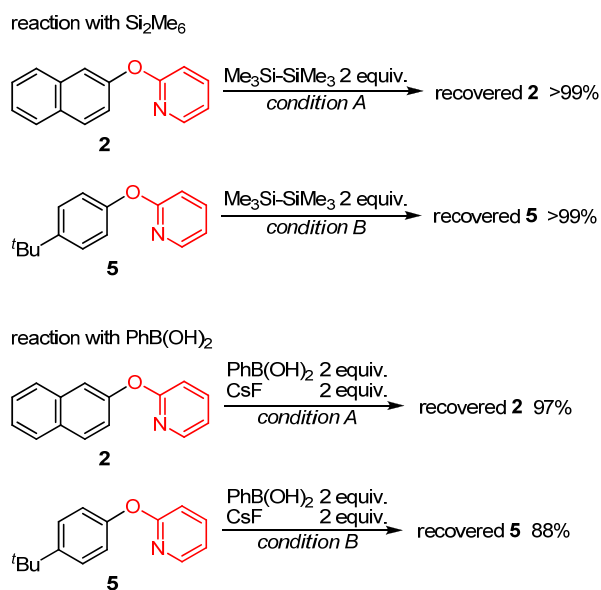
58
59 (Scheme 3 can be found in page 10)

The elementary steps involved in the present rhodium-
catalyzed borylation of aryl 2-pyridyl ethers are outlined
in Scheme 4. The chlororhodium(I) precursor **A** initially
generates borylrhodium(I) species **C** via the sequence of
oxidative addition of B₂(pin)₂/reductive elimination of
ClB(pin).^{19b} Borylrhodium(I) complex **C** subsequently me-
diates the activation of the C(aryl)-O bond in the aryl 2-
pyridyl ether, which should lead to the formation of aryl-
rhodium(I) complex **D**, along with *N*-boryl-2-pyridone **E**
or its tautomer **E'**.³¹ 2-Hydroxypyridine, which can be
formed by the hydrolysis of **E** or **E'**, was observed by ¹H
NMR spectroscopy of the crude reaction mixture (ca.
55%).³² The intermediate **D** eventually forms a borylated
product by the reaction with B₂(pin)₂ with concomitant
regeneration of borylrhodium **C**. The process of the gen-
eration of the catalytically active species (**A**→**B**→**C**) and
the process of product formation and catalyst turnover (**D**
→**F**→**C**) are basically the same as those involved in the
previously reported borylation reaction of nitriles, where
the feasibility of these processes was demonstrated by
computational studies.¹⁹ Accordingly, the key issue is the
mode of action of borylrhodium complex **C** in the C(aryl)-
O bond activation step (**C**→**D**).

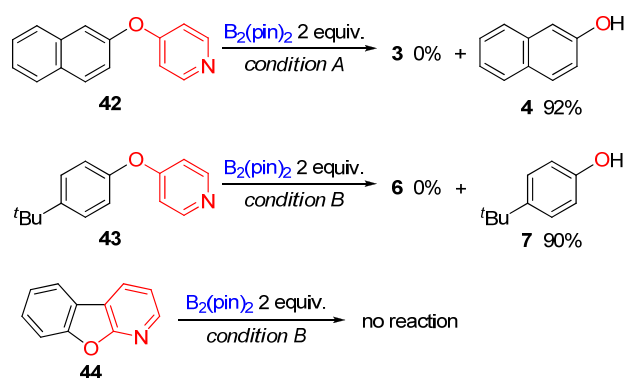
(Scheme 4 can be found in page 11)

To gain insight into the mechanism for C(aryl)-O bond
activation in the present catalytic reaction, several control
experiments were performed. First, the rhodium-
catalyzed reactions of the aryl 2-pyridyl ethers **2** and **5**
were performed with either Si₂Me₆ or PhB(OH)₂ in place
of B₂(pin)₂ to investigate whether the corresponding silyl-
rhodium^{15,22,33} or phenylrhodium³⁴ species can mediate
the activation of a C(aryl)-O bond in a similar manner
(Scheme 5). However, in all cases, none of the C(aryl)-O
bond cleavage products were observed and the starting
material was quantitatively recovered. These results clearly
indicate that the boryl ligand on the rhodium center is
crucial for the activation of aryl 2-pyridyl ethers. It was
also confirmed that the reactions of the corresponding 4-
pyridyl ethers **42** and **43** did not give the C(aryl)-O bond
cleavage products, but rather resulted in cleavage of the
C(4-pyridyl)-O bond to form 2-naphthol and 4-*tert*-
butylphenol, respectively (Scheme 6).³⁵ Moreover, no re-
action occurred with conformationally restricted cyclic
substrate **44** under the current optimized reaction condi-
tions. These results suggest that the presence of a nitro-
gen atom at the appropriate position in the substrate is
essential for the formation of the transition state for the
desired C(aryl)-O bond activation.

Scheme 5. Reaction with Silylrhodium and Phenylrhodium Species



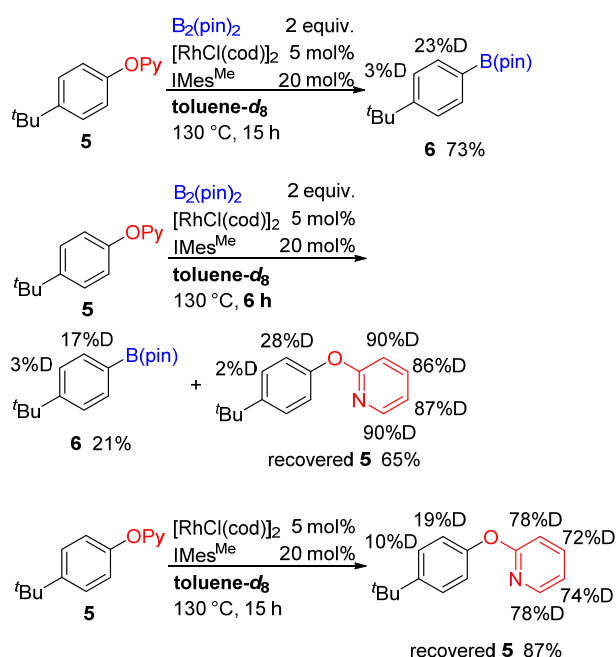
Scheme 6. Control Experiments



Another interesting issue is the selectivity between the activation of an ortho C-H bond and C(aryl)-O bond. Despite a number of reported examples of ortho functionalization of aryl 2-pyridyl ethers using palladium^{11b,11c,11f,11h-k} and ruthenium^{11a,11d,11g} catalysts, no ortho borylated products were observed with all substrates investigated under the current reaction conditions. Interestingly, deuterium incorporation at the ortho position was observed when the reaction was performed in toluene- d_8 .^{36,37} For example, the rhodium-catalyzed reaction of **5** with $\text{B}_2(\text{pin})_2$ in toluene- d_8 was found to involve H/D exchange between the aromatic C-H bonds of **5** and **6** and the deuterated solvent (Scheme 7). At the time of 35% conversion of **5**, deuterium was incorporated at 17% and 28% of the ortho C-H bonds of **6** and the recovered substrate **5**, respectively. This observation indicates that activation of the ortho C-H bond by the rhodium catalyst did occur under the current reaction conditions. However, the rate of C-H activa-

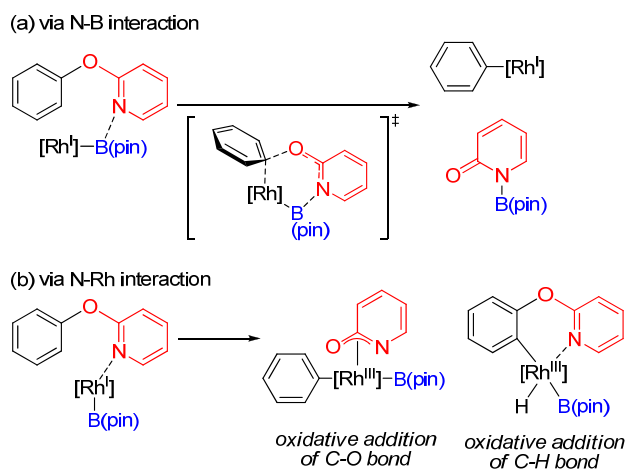
tion was relatively slow compared with that for activation of the C(aryl)-O and C(pyridyl)-H bonds. Importantly, H/D exchange occurred to a similar extent when the reaction was performed in the absence of $\text{B}_2(\text{pin})_2$. Thus, these nonproductive C-H bond activation reactions were not mediated by borylrhodium species **C**, which is responsible for the desired borylation of the C(aryl)-O bond, but rather by a simpler rhodium species, such as $\text{RhCl}(\text{cod})(\text{IMes}^{\text{Me}})$.³⁸

Scheme 7. H/D Exchange with Deuterated Solvent



Based on the above-mentioned control experiments and the observed H/D exchange, we believe that C(aryl)-O bond activation is initiated by the Lewis acid/base interaction between the boryl ligand and the 2-pyridyl moiety (Scheme 8a).³⁹ This interaction not only brings the rhodium center close to the C(aryl)-O bond, but also makes the 2-pyridyloxy group a better leaving group by imparting pyridinium-like character. Collectively, C(aryl)-O bond activation proceeds via a six-membered cyclic transition state, as we initially envisioned. However, we cannot completely exclude an oxidative addition pathway,^{24c,24d} which may be facilitated by the coordination of the pyridine ring to the rhodium center (Scheme 8b). This mode of coordination could also facilitate oxidative addition of an ortho C-H bond, which allows for the formation of a relatively stable six-membered metallacyclic intermediate. However, considering that ortho C-H bond activation is relatively slow compared with C(aryl)-O bond activation under the current reaction conditions, we prefer the possible mechanism shown in Scheme 8a over that shown in Scheme 8b.

Scheme 8. Two Possible Mechanisms for the Cleavage of the Ar-OPy Bond



Conclusions

We have developed a rhodium-catalyzed borylation reaction for aryl 2-pyridyl ethers through the selective cleavage of C(aryl)-OPy bonds. This reaction proceeds under neutral conditions and can be applied to a wide range of substrates that can be easily obtained from the corresponding phenol derivatives. Mechanistic studies revealed that this reaction was specifically catalyzed by a borylrhodium species, and the 2-pyridyl group plays a crucial role in the C(aryl)-O activation step. This method allows for the use of the OPy group as a convertible directing group in C-H bond activation reactions. Further investigation of new catalytic transformations using the unique polarity of borylrhodium(I) species, such as **C**, is currently underway in our laboratory.⁴⁰

ASSOCIATED CONTENT

Supporting Information. Detailed experimental procedures and characterization of new compounds are available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

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Table 3. Rh-Catalyzed Borylation of Aryl Pyridyl Ethers with B₂(pin)₂

$$\text{R-Opy} + \text{B}_2(\text{pin})_2 \xrightarrow[\text{toluene}]{\text{[RhCl(cod)]}_2, \text{ligand}} \text{R-B}(\text{pin})_2$$

2 equiv.

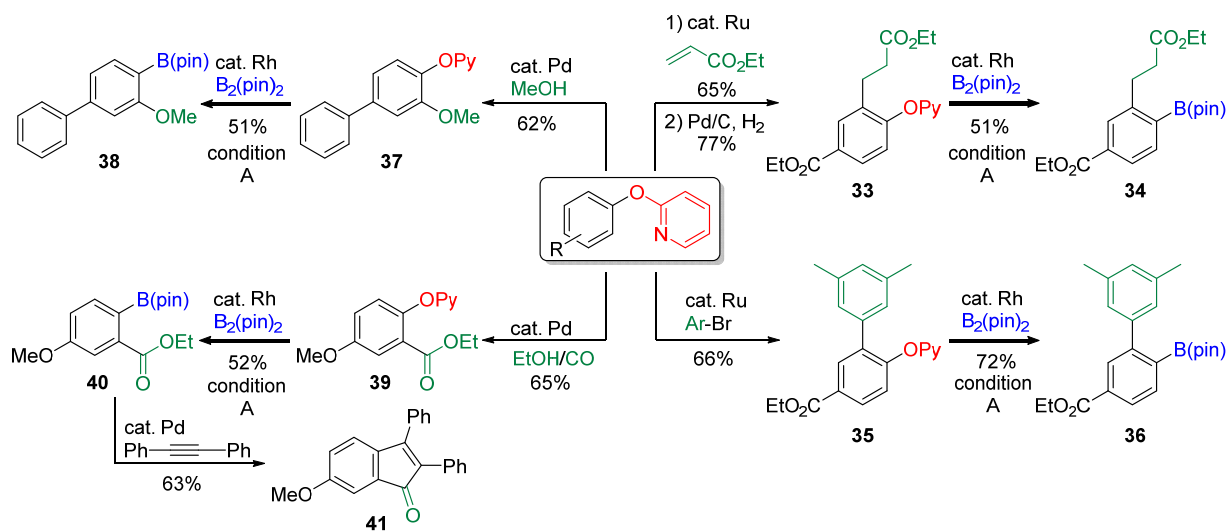
condition A: PCy₃, 100 °C
 condition B: IMes^{Me}, 130 °C
 condition C: IMXy^{Me}, 160 °C

X = Me: IMes^{Me}
 OMe: IMXy^{Me}

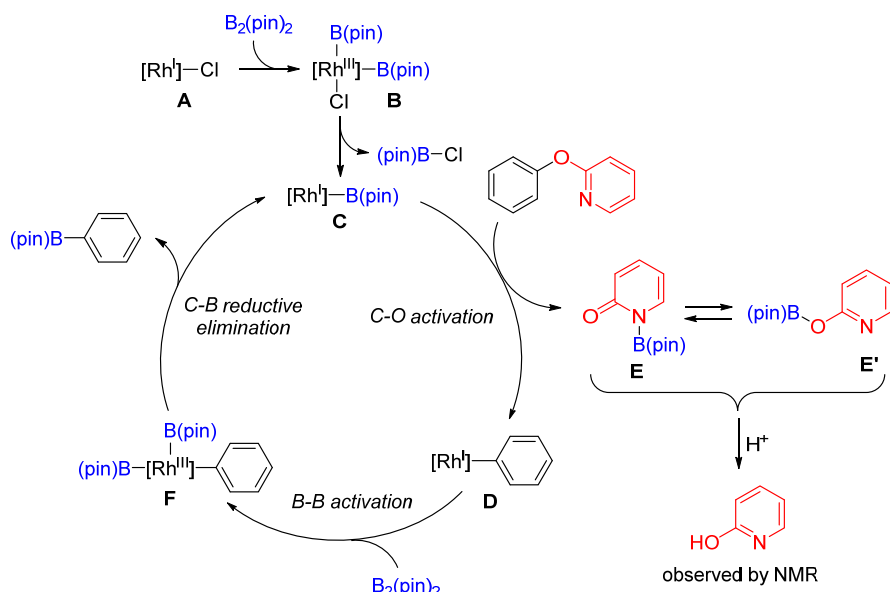
entry	ether	product	condition ^a	yield (%)	entry	ether	product	condition ^a	yield (%)
1			B	77	26			A	89
2			B	80	27			B	59
3			B	68	28			A	61
4			B	71	29			B	65
5			B	62	30			B	82
6			B ^{b,c}	70	31			B	60
7			A	21 ^d	32			B	60
8			B	0	33			B ^{b,c}	65
9			A ^e	65	34			B	66
10			B	61	35			A	68
11			A ^e	75	36			B	21
12			B	30	37			A	62
13			A ^{b,e}	72	38			B	21
14			B	0	39			B	68
15			A ^{b,e}	66	40			A	68
16			B	0	41			B	21
17			B	68	42			A	62
18			B ^f	42	43			B	21
19			C	63	44			A	62
20			B ^f	50	45			B	21
21			C	70	46			A	62
22			B	40	47			B	21
23			A ^{c,e}	53	48			A	62
24			B	30	49			B	21
25			B	60	50			A	62
			B	60	51			B	21
			B	60	52			A	62
			B	60	53			B	21
			B	60	54			A	62
			B	60	55			B	21

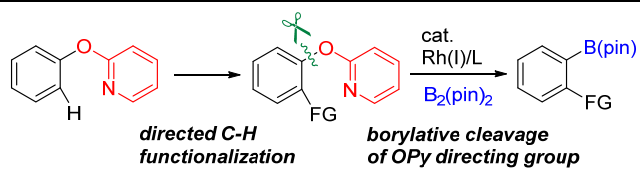
^a Condition A: substrate (0.50 mmol), B₂(pin)₂ (1.0 mmol), [RhCl(cod)]₂ (0.025 mmol), PCy₃ (0.15 mmol), toluene (0.50 mL) for 15 h at 100 °C. Condition B: substrate (0.50 mmol), B₂(pin)₂ (1.0 mmol), [RhCl(cod)]₂ (0.025 mmol), IMes^{Me} (0.10 mmol), toluene (0.50 mL) for 15 h at 130 °C. Condition C: substrate (0.50 mmol), B₂(pin)₂ (1.5 mmol), [RhCl(cod)]₂ (0.050 mmol), ligand (0.20 mmol), toluene (0.50 mL) for 15 h at 160 °C. ^b 3 equiv of B₂(pin)₂ was used. ^c Run for 48 h. ^d 71% of substrate was recovered. ^e Run at 120 °C. ^f Run at 160 °C.

Scheme 3. Sequential Functionalization of Ortho C-H Bond/Borylation of Pyridyl Ethers



Scheme 4. Plausible Catalytic Cycle





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